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FULL ESTIMATED COST

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=> s futhan/cn

L1 1 FUTHAN/CN

=> d

L1 ANSWER 1 OF 1 REGISTRY COPYRIGHT 2000 ACS

RN 82956-11-4 REGISTRY

CN Benzoic acid, 4-[(aminoiminomethyl)amino]-, 6-(aminoiminomethyl)-2naphthalenyl ester, dimethanesulfonate (9CI) (CA INDEX NAME)

OTHER NAMES:

CN 6-Amidino-2-naphthyl p-guanidinobenzoate

CN FUT 175

CN Futhan

CN Nafamostat mesilate

CN Nafamostat mesylate

MF C19 H17 N5 O2 . 2 C H4 O3 S

LC STN Files: ANABSTR, BEILSTEIN\*, BIOBUSINESS, BIOSIS, CA, CAPLUS, CBNB, CIN, DDFU, DRUGPAT, DRUGU, DRUGUPDATES, EMBASE, IMSDIRECTORY, IPA, MRCK\*, PHAR, PROMT, RTECS\*, TOXLINE, TOXLIT, USAN, USPATFULL (\*File contains numerically searchable property data)

CM 1

CRN 81525-10-2 CMF C19 H17 N5 O2

CM 2

CRN 75-75-2 CMF C H4 O3 S

204 REFERENCES IN FILE CA (1967 TO DATE)
2 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA
206 REFERENCES IN FILE CAPLUS (1967 TO DATE)

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=> file caplus, uspa ll, biosis, medline
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CA INDEXING COPYRIGHT (C) 2000 AMERICAN CHEMICAL SOCIETY (ACS)
FILE 'BIOSIS' ENTERED AT 10:48:29 ON 07 MAR 2000
COPYRIGHT (C) 2000 BIOSIS(R)
FILE 'MEDLINE' ENTERED AT 10:48:29 ON 07 MAR 2000
=> s 11
L2
           338 L1
=> s ?cardio? or ?heart?
       1915003 ?CARDIO? OR ?HEART?
L3
=> s 12 and 13
            48 L2 AND L3
T.4
=> s 12 (P) 13
L5
             5 L2 (P) L3
=> dup rem 14
PROCESSING COMPLETED FOR L4
L6
             45 DUP REM L4 (3 DUPLICATES REMOVED)
=> d 1-5 bib, ab 15
L5
     ANSWER 1 OF 5 CAPLUS COPYRIGHT 2000 ACS
ΑN
     1998:12232 CAPLUS
     128:86085
DN
     Elimination of Kupffer cells and administration of protease inhibitor
ΤI
     improve graft viability and prevent reperfusion injury in NHBD
ΑU
     Tsukamoto, S.; Ohkohchi, N.; Orii, T.; Fukumóri, T.; Asakura, T.;
     Takayama, J.; Kato, H.; Satomi, S.
     Second Dep. Surgery, Tohoku Univ. School Medicine, Sendai, Japan Transplant. Proc. (1997), 29(8), 3463-3464
CS
SO
     CODEN: TRPPA8; ISSN: 0041-1345
PΒ
     Elsevier Science Inc.
     Journal
DT
LA
     English
     The aim of this study was to det. Whether liver grafts from
AΒ
     non-heartbeating donors are suitable for clin. liver transplantation.
The
     authors report that the elimination of Kupffer cells and administration
of
     NM (nafamostat mesilate), a serine protease inhibitor that inhibits
    phospholipase A2, improves graft viability and prevents reperfusion
injury
```

in non-heartbeating donors.

ANSWER 2 OF 5 LUS COPYRIGHT 2000 ACS 1998:183 CAPLUS 128:110669 TIEffects of protease inhibitors on postischemic recovery of the heart Shibata, Toshihiko; Yamamoto, Fumio; Suehiro, Shigefumi; Kinoshita, CS Second Dep. of Surgery, Osaka City University Medical School, Osaka, 545, SO Cardiovasc. Drugs Ther. (1997), 11(4), 547-556 CODEN: CDTHET; ISSN: 0920-3206 PΒ Kluwer Academic Publishers DTJournal LA English AΒ It is well known that activation of proteases in the lysosomes and cytosol is one of the mechanisms of ischemic injury. It might thus be beneficial to det. whether the addm. of several clin. available protease inhibitors to a cardioplegic soln. can improve its protective ability. Using an isolated working rat heart prepn., the effects of several protease inhibitors (serine protease inhibitors; nafamostat mesylate and gabexate mesylate, a thiol-protease inhibitor; NCO 700; and a urinary trypsin inhibitor, urinastatin) on the postischemic recovery of function and enzyme leakage were investigated in this study. These protease inhibitors were added to either the cardioplegic soln. or reperfusion soln. The addn. of each of the protease inhibitors, except urinastatin, to the cardioplegic soln. improved the postischemic recovery of function and reduced enzyme leakage. The dose-response characteristics of these three protease inhibitors were bell shaped, and the optimal concns. of nafamostat mesylate, gabexate mesylate, and NCO-700 were 5 .mu.M, 100 .mu.M, and 20 .mu.M, resp. In contrast to the results of the preischemic treatment study, the addn. of any of the protease inhibitors to the perfusion medium during Langendorff reperfusion failed to improve the postischemic recovery of function and to reduce enzyme leakage. Surprisingly, the addn. of NCO-700 to the reperfusion soln. at a concn. of 5 .mu.M or higher had rather harmful effects on both functional recovery and enzyme leakage. These findings suggest that serine and thiol proteases may play an important role in myocardial injury during ischemia, but not necessarily during reperfusion. L5 ANSWER 3 OF 5 CAPLUS COPYRIGHT 2000 ACS 1997:31579 ÇAPLUS ΑN DN 126:84283 The effect and pharmacokinetics of nafamostat mesilate adjunct to cold ΤI nondepolarizing cardioplegia in a canine model of cardiac preservation Sunamori, Makoto; Yoshida, Tetsuya; Miyamoto, Hisashi; Wang, Yigang; ΑU Suzuki, Akio CS School Medicine, Tokyo Medical and Dental University, Tokyo, 113, Japan SO Transplant Int. (1996), 9(4), 364-369 CODEN: TRINE5; ISSN: 0934-0874 PB Springer DT Journal English LA AΒ The effects of nafamostat mesilate (NM) on myocardial, biochem., and functional changes in canine hearts were examd. An isolated heart was preserved for 6 h at 5.degree. and then reperfused for 2 h at 37.degree..

LA English

The effects of nafamostat mesilate (NM) on myocardial, biochem., and functional changes in canine hearts were examd. An isolated heart was preserved for 6 h at 5.degree. and then reperfused for 2 h at 37.degree. NM was added to the cardioplegic soln. At both 10-7M and 10-6M, NM was able to maintain myocardial cAMP at a normal level and to reduce cGMP concns. at the end of both preservation and reperfusion. The serum N-acetyl-.beta.-D-glucosaminidase concn. during reperfusion was lower in hearts treated with NM 10-6 or 10-7M NM than in those without NM. Although NM failed to preserve myocardial concns. of adenine nucleotide compds., NM at 10-7M maintained the .+-. dp/dt of the left ventricle after

reperfusion at same level as in the nonische control group and better than NM alo-6M or no NM. Myocardial uptake of 10-5M NM was 55% during the 6-h preservation and 29% during the 2-h reperfusion. It is concluded that addn. of 10-7M NM to a nondepolarizing soln. does not preserve myocardial adenine nucleotide concns. but does facilitate the recovery of left ventricular function. NM at 10-5M seems to have a high affinity for the myocardium and may depress the recovery of left ventricular function.

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L5 ANSWER 4 OF 5 CAPLUS COPYRIGHT 2000 ACS
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AN 1993:404342 CAPLUS

DN 119:4342

TI Efficacy of futhan rinse solution following rat heart preservation AU Urushihara, Takashi; Sumimoto, Kazuo; Sumimoto, Ryo; Ikeda, Masanobu;

Fukuda, Yasuhiko; Dohi, Kiyohiko

CS Sch. Med., Hiroshima Univ., Hiroshima, Japan

SO Nippon Geka Gakkai Zasshi (1992), 93(12), 1514

CODEN: NGGZAK; ISSN: 0301-4894

DT Journal

LA Japanese

AB The efficacy of futhan rinse soln. following rat heart preservation was studied and compared with that of physiol. saline soln. and Carolina Rinse-II soln. The graft survival after 18-h preservation was 100, 50, and 50% in futhan rinse, physiol. saline, and Carolina Rinse-II group, resp. These results suggest the potential of clin. application of futhan rinse soln.

L5 ANSWER 5 OF 5 CAPLUS COPYRIGHT 2000 ACS

AN 1989:546489 CAPLUS

DN 111:146489

 ${
m TI}$  Experimental study on the usefulness of the protease inhibitor, nafamostat

mesilate (FUT), as an anticoagulant in left heart bypass

AU Saito, A.; Moro, H.; Eguchi, S.; Yokosawa, T.

CS Sch. Med., Niigata Univ., Niigata, Japan

SO Jinko Zoki (1989), 18(2), 453-6 CODEN: JNZKA7; ISSN: 0300-0818

DT Journal

LA Japanese

AB The left heart bypass was performed in seven mongrel dogs for 4 h using a protease inhibitor nafamostat mesilate (FUT) as an anticoagulant. Bypass flow was 200 mL/min (20% of cardiac output) and FUT was infused continuously via the inflow cannula. ACT (activated clotting time), platelet counts, activated partial thrombin time and fructose 1,6-diphosphate were measured during the bypass. The results show that nafamostat mesilate is a useful and safe anticoagulant for mech. circulatory support.

=> d 16 1-45 bib

=>

L6 ANSWER 1 OF 45 CAPLUS COPYRIGHT 2000 ACS

AN 1999:595348 CAPLUS

DN 131:225828

TI Methods of diagnosis and triage using cell activation measures

IN Stoughton, Roland B.; Schmid-Schonbein, Geert W.; Hugli, Tony E.; Kistler,

Erik

PA Cell Activation, Inc., USA; The Regents of the University of California; The Scripps Research Institute

SO PCT Int. Appl., 184 pp. CODEN: PIXXD2

DT Patent

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FAN.CNT 1
     PATENT NO.
                   KIND DATE
                                            APPLICATION NO. DATE
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                             _____
                                             -----
ΡI
     WO 9946367
                       A2
                             19990916
                                             WO 1999-US5247
                                                               19990311
     WO 9946367
                       A3
                             19991209
         W: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ,
              DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS,
              JP, KE, KG, KP, KR, KZ, LC, LR, LS, LT, LU, LV, MD, MG, MK, MN,
             MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM,
             TR, TT, UA, UG, US, UZ, VN, YU, ZW, AM, AZ, BY, KG, KZ, MD, RU,
             TJ, TM
         RW: GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW, AT, BE, CH, CY, DE, DK,
             ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG,
             CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                       A1 19990927
     AU 9931829
                                           AU 1999-31829
                                                               19990311
PRAI US 1998-38894
                       19980311
     WO 1999-US5247
                       19990311
     ANSWER 2 OF 45 BIOSIS COPYRIGHT 2000 BIOSIS
L6
     1999:200615 BIOSIS
ΑN
DN
     PREV199900200615
ΤI
     Prevention of neointimal formation by a serine protease inhibitor,
     FUT-175, after carotid balloon injury in rats.
     Sawada, Motoshi; Yanamoto, Hiroji (1); Nagata, Izumi; Hashimoto, Nobuo; Nakahara, Ichiro; Akiyama, Yoshinori; Kikuchi, Haruhiko
ΑU
     (1) Laboratory for Cerebrovascular Disorders, National Cardio-Vascular
CS
     Center Research Institute, 5-7-1 Fujishiro-dai, Suita, 565-8565 Japan Stroke, (March, 1999) Vol. 30, No. 3, pp. 644-650.
SO
     ISSN: 0039-2499.
\mathsf{DT}
     Article
LA
     English
L6
     ANSWER 3 OF 45 BIOSIS COPYRIGHT 2000 BIOSIS
     1999:470874 BIOSIS
ΑN
DN
     PREV199900470874
TI
     Direct evidence of the role of mucosal mast cell activation in the
     pathogenesis of intestinal ischemia-reperfusion injury in rats.
     Kimura, T. (1); Andoh, A. (1); Fukuda, M. (1); Tsujikawa, T. (1); Sasaki,
ΑIJ
     M. (1); Fujiyama, Y. (1); Bamba, T. (1)
CS
     (1) Dept. of Internal Medicine, Shiga University of Medical Science, Otsu
     Japan
     Journal of Parenteral and Enteral Nutrition, (Sept. Oct., 1999) Vol. 23,
SO
     No. 5 SUPPL., pp. S149.
     Meeting Info.: International Symposium on Growth Factors and Nutrients in
     Intestinal Health and Disease Osaka, Japan October 31-November 3, 1998
     ISSN: 0148-6071.
DT
     Conference
LA
     English
     ANSWER 4 OF 45 BIOSIS COPYRIGHT 2000 BIOSIS
L6
AN
     1998:475040 BIOSIS
     PREV199800475040
DN
     Interleukin-6 derived from hypoxic myocytes promotes neutrophil-mediated
TΙ
     reperfusion injury in myocardium.
     Sawa, Yoshiki (1); Ichikawa, Hajime; Kagisaki, Koji; Ohata, Toshihiro;
ΑU
     Matsuda, Hikaru
     (1) First Dep. Surg., Osaka Univ. Med. Sch., 2-2 Yamada-oka, Suita, Osaka
CS
     565 Japan
     Journal of Thoracic and Cardiovascular Surgery, (Sept., 1998) Vol. 116,
SO
     No. 3, pp. 511-517.
     ISSN: 0022-5223.
DT
     Article
LA
     English
```

ANSWER 5 OF 45 CAPLUS COPYRIGHT 2000 ACS

English

LA

L6

```
ΑN
      1998:12232 CAP
 DN
      128:86085
 TΙ
      Elimination of Kupffer cells and administration of protease inhibitor
      improve graft viability and prevent reperfusion injury in NHBD
 ΑU
      Tsukamoto, S.; Ohkohchi, N.; Orii, T.; Fukumori, T.; Asakura, T.;
      Takayama, J.; Kato, H.; Satomi, S.
 CS
      Second Dep. Surgery, Tohoku Univ. School Medicine, Sendai, Japan
      Transplant. Proc. (1997), 29(8), 3463-3464
 SO
      CODEN: TRPPA8; ISSN: 0041-1345
 PB
      Elsevier Science Inc.
 \mathsf{DT}
      Journal
 LA
      English
 L6
      ANSWER 6 OF 45 CAPLUS COPYRIGHT 2000 ACS
 ΑN
      1998:183 CAPLUS
 DN
      128:110669
     Effects of protease inhibitors on postischemic recovery of the
 TI
ΑU
     Shibata, Toshihiko; Yamamoto, Fumio; Suehiro, Shigefumi; Kinoshita,
     Second Dep. of Surgery, Osaka City University Medical School, Osaka, 545,
 CS
 SO
     Cardiovasc. Drugs Ther. (1997), 11(4), 547-556
     CODEN: CDTHET; ISSN: 0920-3206
PB
     Kluwer Academic Publishers
DT
     Journal
     English
LA
     ANSWER 7 OF 45 USPATFULL
L6
ΑN
       96:94448 USPATFULL
ΤI
       Perfusion and storage solution containing sodium lactobionate, sodium
       dihydrogenphosphate, raffinose, glutathione, allopurinol and nafamostat
       mesylate
IN
       Dohi, Kiyohiko, Hiroshima, Japan
       Urushihara, Takashi, Hiroshima, Japan
       Iwata, Masanori, Chiba, Japan
       Torii Pharmaceutical Co., Ltd., Tokyo, Japan (non-U.S. corporation)
PΑ
PΙ
       US 5565317 19961015
       WO 9400008 19940106
       US 1994-347490 19941206 (8)
ΑI
       WO 1993-JP219 19930223
              19941206 PCT 371 date
19941206 PCT 102(e) date
PRAI
       JP 1992-168977
                           19920626
DT
       Utility
       Primary Examiner: Naff, David M.; Assistant Examiner: Saucier, S.
EXNAM
LREP
       Beveridge, DeGrandi, Weilacher & Young, L.L.P.
       Number of Claims: 17
CLMN
ECL
       Exemplary Claim: 1
DRWN
       No Drawings
LN.CNT 263
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
L6
     ANSWER 8 OF 45 BIOSIS COPYRIGHT 2000 BIOSIS
     1996:321861 BIOSIS
ΑN
DN
     PREV199699044217
ΤI
     Pharmacological therapeutic prospects of cerebral vasospasm.
ΑU
     Serv. Univ. d'Anesthesie Reanim., CHR de la Citadelle, 4000 Liege Belgium
CS
SO
     Annales Francaises d'Anesthesie et de Reanimation, (1996) Vol. 15, No. 3,
     pp. 374-381.
     ISSN: 0750-7658.
DT
     General Review
```

French

French; English

LA SL

- L6 ANSWER 9 OF 45 LUS COPYRIGHT 2000 ACS ΑN 1997:31579 CAPL DN 126:84283 The effect and pharmacokinetics of nafamostat mesilate adjunct to cold ΤI nondepolarizing cardioplegia in a canine model of cardiac ΑU Sunamori, Makoto; Yoshida, Tetsuya; Miyamoto, Hisashi; Wang, Yigang; Suzuki, Akio School Medicine, Tokyo Medical and Dental University, Tokyo, 113, Japan CS SO Transplant Int. (1996), 9(4), 364-369 CODEN: TRINE5; ISSN: 0934-0874 PΒ Springer DTJournal LAEnglish L6 ANSWER 10 OF 45 BIOSIS COPYRIGHT 2000 BIOSIS ΑN 1996:408501 BIOSIS DN PREV199699130857 TΙ Effect of FUT-175 on postoperative organ dysfunction following open heart surgery with cardiopulmonary bypass (with special reference to soluble adhesion molecules. Shimamoto, A. (1); Sato, T.; Kondo, C.; Shomura, Y. (1); Hioki, I. (1); ΑU Tempaku, H. (1); Maze, Y. (1); Takao, M. (1); Onoda, K.; Tani, K.; K.; Shimpo, H.; Yada, I. (1) Dep. Thorac. and CV Surg., Mie Univ. Sch. Med., 2-174 Edobashi, Tsu, Mie 514 Japan Japanese Journal of Artificial Organs, (1996) Vol. 25, No. 2, pp. 361-364. ISSN: 0300-0818. DTArticle LA Japanese SLJapanese; English L6 ANSWER 11 OF 45 BIOSIS COPYRIGHT 2000 BIOSIS ΑN 1997:33421 BIOSIS DN PREV199799339824 ΤI Investigation of the anti-complement agents, FUT-175 and K76COOH, in discordant xenotransplantation. Kobayashi, T.; Neethling, F. A.; Taniguchi, S.; Ye, Y.; Niekrasz, M.; ΑU Koren, E.; Hancock, W. W.; Takagi, H.; Cooper, D. K. C. (1) (1) Transplantation Biology Res. Cent., Mass. General Hosp., MGH-East, CS Boston, MA 02129 USA Xenotransplantation, (1996) Vol. 3, No. 3, pp. 237-245. SO ISSN: 0908-665X. DTArticle English LA L6 ANSWER 12 OF 45 CAPLUS COPYRIGHT 2000 ACS ΑN 1996:105043 CAPLUS DN 124:193852 Nafamostat mesylate, a broad spectrum protease inhibitor, modulates TΙ platelet, neutrophil and contact activation in simulated extracorporeal circulation ΑU Sundaram, Sumuk; Gikakis, Nicolas; Hack, C. Erik; Niewiarowski, Stefan; Edmunds, L. H., Jr.; Rao, A. Koneti; Sun, Ling; Cooper, S. L.; Colman,
- Robert W.
- Sol Sherry Thrombosis Res. Cent., Temple Univ. Sch. Med., Philadelphia, PA, 19140, USA
- SO Thromb. Haemostasis (1996), 75(1), 76-82 CODEN: THHADQ; ISSN: 0340-6245
- DT Journal
- LA English
- L6 ANSWER 13 OF 45 BIOSIS COPYRIGHT 2000 BIOSIS
- 1996:112706 BIOSIS ΑN

DN PREV19969868484 ΤI Attenuation of diopulmonary bypass-derived inflammatory reactions reduces myocardial reperfusion injury in cardiac operations. ΑU Sawa, Yoshiki; Shimazaki, Yasuhisa; Kadoba, Keishi; Masai, Takashi; Fukuda, Hirotsugu; Ohata, Toshihiro; Taniguchi, Kazuhiro; Matsuda, Hikaru CS (1) First Dep. Surgery, Osaka Univ. Med. Sch., 2-2 Yamada-oka, Suita, Osaka 565 Japan SO Journal of Thoracic and Cardiovascular Surgery, (1996) Vol. 111, No. 1, pp. 29-35. ISSN: 0022-5223. DT Article LAEnglish L6 ANSWER 14 OF 45 USPATFULL ΑN 95:45354 USPATFULL TIThermoplastic polymer composition and medical devices made of the same ΙN Endo, Fumiaki, Fuji, Japan Saiga, Nobuko, Hadano, Japan PΑ Terumo Kabushiki Kaisha, Tokyo, Japan (non-U.S. corporation) PΙ US 5417981 19950523 US 1993-53499 19930428 (8) ΑТ PRAI JP 1992-136329 19920428 Utility Primary Examiner: Kight, III, John; Assistant Examiner: Dodson, Shelley EXNAM LREP Burns, Doane, Swecker & Mathis CLMN Number of Claims: 24 ECL Exemplary Claim: 1 No Drawings DRWN LN.CNT 489 CAS INDEXING IS AVAILABLE FOR THIS PATENT. 1.6 ANSWER 15 OF 45 BIOSIS COPYRIGHT 2000 BIOSIS ΑN 1996:142434 BIOSIS DN PREV199698714569 TΙ Nafamostat mesylate is the first choice anticoagulant for continuous renal replacement therapy. ΑU Sugai, Takao; Hirasawa, Hiroyuki; Ohtake, Yoshio; Oda, Shigeto; Nakanishi, Kazuya; Kitamura, Nobuya; Matsuda, Kenichi; Kawabe, Touichi; Ueno, Hirokazu; Touma, Takayuki; Yokohari, Kenji Dep. Emergency, Critical Care Med., Chiba Univ. Sch. Med., Chiba Japan Blood Purification, (1995) Vol. 13, No. 6, pp. 388. CS SO Meeting Info.: International Conference on Continuous Renal Replacement Therapies San Diego, California, USA November 8-10, 1995 ISSN: 0253-5068. DT Conference LA English L6 ANSWER 16 OF 45 BIOSIS COPYRIGHT 2000 BIOSIS ΑN 1995:243942 BIOSIS DN PREV199598258242 TΙ The anti-complement effects of FUT-175 on myocardial ischemia/reperfusion injury in the blood-perfused isolated rabbit hearts. ΑU Yokota, Syunji (1); Kan-No, Satoshi; Saitoh, Yoshiaki; Kasama, Kikuko; Ohara, Naoki; Ono, Hiroshi CS (1) Lab. Applied Pharmacology, Hatano Res. Inst., Food Drug Safety Center, Kanagawa 257 Japan SO Japanese Journal of Pharmacology, (1995) Vol. 67, No. SUPPL. 1, pp. 280P. Meeting Info.: 68th Annual Meeting of the Japanese Pharmacological Society Nagoya, Japan March 25-28, 1995

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ISSN: 0021-5198
 DT
      Conference
 LA
      English
 L6
      ANSWER 17 OF 45 BIOSIS COPYRIGHT 2000 BIOSIS
 AN
      1995:257247 BIOSIS
 DN
      PREV199598271547
      Inhibitory effect of FUT-175 on the production of interleukin 8 and
 ΤI
      polymorphonuclear leukocyte elastase.
      Kikuchi, Mitsuru (1); Endo, Shigeatsu; Inada, Katsuya; Yamashita,
 ΑU
      Hisahiko; Takauwa, Tetsuya; Nakae, Hajime; Kasai, Takeshi; Baba, Nobuo;
      Yamada, Yasuhiko
 CS
      (1) Critical Care Emergency Cent., Iwate Med. Univ., 19-1 Uchimaru,
      Morioka 020 Japan
 SO
      Research Communications in Molecular Pathology and Pharmacology, (1995)
      Vol. 87, No. 3, pp. 269-274.
 DТ
      Article
 LA
      English
      ANSWER 18 OF 45 CAPLUS COPYRIGHT 2000 ACS
 L6
 ΑN
      1994:129080 CAPLUS
 DN
      120:129080
 TI
      Organ-preserving fluid
      Dohi, Kiyohiko; Urushihara, Takashi; Iwata, Masanori
 ΙN
      Torii and Co., Ltd., Japan
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 SO
      PCT Int. Appl., 13 pp.
      CODEN: PIXXD2
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 FAN.CNT 1
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                                            APPLICATION NO.
                                                              DATE
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                             19940106
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         RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE
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PRAI JP 1992-168977
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L6
     ANSWER 19 OF 45 USPATFULL
AN
       94:71052 USPATFULL
ΤI
       Gas permeable thrombo-resistant coatings and methods of manufacture
ΤN
       Winters, Suzanne, Salt Lake City, UT, United States
       Solen, Kenneth A., Orem, UT, United States
       Sanders, Clifton G., Salt Lake City, UT, United States
       Mortensen, JD, Sandy, UT, United States
       Berry, Gaylord, Salt Lake City, UT, United States
PΑ
       Cardiopulmonics, Inc., Salt Lake City, UT, United States (U.S.
       corporation)
PΙ
       US 5338770 19940816
ΑI
       US 1990-509063 19900412 (7)
DCD
       20101116
RLI
       Continuation-in-part of Ser. No. US 1988-215014, filed on 5 Jul 1988,
       now patented, Pat. No. US 5262451 which is a continuation-in-part of
       Ser. No. US 1988-204115, filed on 8 Jun 1988, now patented, Pat. No. US
       4850958
DT
       Utility
EXNAM
       Primary Examiner: Szekely, Peter
LREP
       Workman Nydegger Jensen
CLMN
       Number of Claims: 33
ECL
       Exemplary Claim: 1
```

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

- L6 ANSWER 20 OF 45 BIOSIS COPYRIGHT 2000 BIOSIS
- AN 1995:67714 BIOSIS
- DN PREV199598082014
- TI Platelet protection with FUT-175 during cardiopulmonary bypass surgery: Electron microscopic study.
- AU Takagi, K.; Kondo, C.; Tanaka, K.; Yada, I.; Kusagawa, M.
- CS Dep. Thorac. Surg., Mie Univ. Sch. Med., 2-174 Edobashi, Tsu, Mie 514 Japan
- SO Japanese Journal of Artificial Organs, (1994) Vol. 23, No. 5, pp. 1089-1094.
  ISSN: 0300-0818.
- DT Article
- LA Japanese
- SL Japanese; English
- L6 ANSWER 21 OF 45 BIOSIS COPYRIGHT 2000 BIOSIS
- AN 1994:384344 BIOSIS
- DN PREV199497397344
- TI Cardiac xenotransplantation from pig to Japanese monkey with splenectomy, tacrolims, filtration plasmapheresis, and nafamstat mesilate.
- AU Kawauchi, M. (1); Takeda, M.; Nakajima, J.; Matsumoto, J.; Furuse, A.
- CS (1) Dep. Thoracic Surg., Univ. Tokyo, 7-3-1 Hongo, Bunkyo-ku, Tokyo 113 Japan
- Transplantation Proceedings, (1994) Vol. 26, No. 3, pp. 1076-1077. Meeting Info.: Second International Congress on Xenotransplantation Cambridge, England, UK September 26-29, 1993 ISSN: 0041-1345.
- DT Conference
- LA English
- L6 ANSWER 22 OF 45 BIOSIS COPYRIGHT 2000 BIOSIS
- AN 1995:24486 BIOSIS
- DN PREV199598038786
- TI Effects of protease inhibitor and immunosuppressant on cerebral vasospasm after subarachnoid hemorrhage in rabbits.
- AU Yanamoto, Hiroji; Kikuchi, Haruhiko (1); Okamoto, Shinichiro
- CS (1) Dep. Neurosurg., Kyoto Univ. Med. Sch., Kawahara 54 Syogoin, Sakyo, Kyoto 606 Japan
- SO Surgical Neurology, (1994) Vol. 42, No. 5, pp. 382-387. ISSN: 0090-3019.
- DT Article
- LA English
- L6 ANSWER 23 OF 45 BIOSIS COPYRIGHT 2000 BIOSIS
- AN 1995:24485 BIOSIS
- DN PREV199598038785
- TI Cerebral vasospasm caused by cisternal injection of polystyrene latex beads in rabbits is inhibited by a serine protease inhibitor.
- AU Yanamoto, Hiroji; Kikuchi, Haruhiko (1); Okamoto, Shinichiro; Nozaki, Kazuhiko
- CS (1) Dep. Neurosurg., Kyoto Univ. Med. Sch., Kawahara 54, Syogoin, Sakyo, Kyoto 606 Japan
- SO Surgical Neurology, (1994) Vol. 42, No. 5, pp. 374-381. ISSN: 0090-3019.
- DT Article
- LA English
- L6 ANSWER 24 OF 45 BIOSIS COPYRIGHT 2000 BIOSIS
- AN 1994:191784 BIOSIS
- DN PREV199497204784
- TI Effect of nafamostat mesilate on Bradykinin generation and hemodynamic changes during LDL apheresis.

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Kojima, S. (1); ba-Harada, M.; Nomura, S.; Kulinchi, M.; Yamamoto, A. (1) Tohsei Natik Hosp. Japan Artificial Organs, (1994) Vol. 18, No. 2, pp. 138.
 CS
 SO
      Meeting Info.: 5th International Congress of the World Apheresis
       Association Houston, Texas, USA March 9-12, 1994
       ISSN: 0160-564X.
 DT
       Conference
 LA
      English
 Ĺ6
      ANSWER 25 OF 45 BIOSIS COPYRIGHT 2000 BIOSIS
      1994:293247 BIOSIS
 ΑN
 DN
      PREV199497306247
      Effect of FUT-175 (nafamostat mesilate) on the development of shock.
 TI
      Maekawa, Yuriko; Koshiyama, Yoshiko; Kashiwabara, Sanae; Oda, Minoru;
 ΑU
      Iwaki, Masahiro
 CS
      Res. Lab., Torii and Co. Ltd., Chiba 267 Japan
      Japanese Journal of Pharmacology, (1994) Vol. 64, No. SUPPL. 1, pp. 97P.
 SO
      Meeting Info.: 67th Annual Meeting of the Japanese Pharmacological
 Society
      Kyoto, Japan March 21-24, 1994
      ISSN: 0021-5198.
 DT
      Conference
 LA
      English
 L6
      ANSWER 26 OF 45 BIOSIS COPYRIGHT 2000 BIOSIS
 ΑN
      1993:533727 BIOSIS
 DN
      PREV199345120821
      Nafamostat mesilate reduces postoperative blood loss in open heart
 TΙ
      surgery.
      Murase, M.; Maeda, M.; Teranishi, K.; Sakurai, H.; Nishizawa, T.; Koyama,
 ΑU
      T.; Itoh, T.
      Ohgaki Municipal Hosp., Dep. thoracic Surg., 4-86 Minaminokawa, Ohggaki,
 CS
      Gifu 503 Japan
      Japanese Journal of Artificial Organs, (1993) Vol. 22, No. 3, pp.
 SO
 943-946.
     Meeting Info.: Thirtieth Meeting of the Japanese Society for Artificial
      Organs
      ISSN: 0300-0818.
DT
      Article
LA
      Japanese
SL
     Japanese; English
L6
     ANSWER 27 OF 45 CAPLUS COPYRIGHT 2000 ACS
                                                          DUPLICATE 1
ΑN
     1993:508660 CAPLUS
     119:108660
DN
     Prolonging discordant xenograft survival with anticomplement reagents
TT
     K76COOH and FUT175
     Miyagawa, Shuji; Shirakura, Ryota; Matsumiya, Goro; Fukushima, Norihide;
ΑU
     Nakata, Seizoh; Matsuda, Hikaru; Matsumoto, Misako; Kitamura, Hajime;
     Seya, Tsukasa
CS
     Med. Sch., Osaka Univ., Osaka, 553, Japan
SO
     Transplantation (1993), 55(4), 709-13
     CODEN: TRPLAU; ISSN: 0041-1337
DT
     Journal
LA
     English
L6
     ANSWER 28 OF 45 BIOSIS COPYRIGHT 2000 BIOSIS
AN
     1993:392414 BIOSIS
DN
     PREV199396067714
     Effect of nafamostat mesilate on sodium and potassium transport
properties
     in the rabbit cortical collecting duct.
ΑU
     Muto, Shigeaki (1); Imai, Masashi; Asano, Yasushi
CS
     (1) Dep. Nephrol. Pharmacol., Jichi Med. Sch., 3311-1 Minamikawachi,
     Tochigi 329-04 Japan
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British Journal of Pharmacology, (1993) Vol. 109, No. 3, pp. 673-678.

SO

- ISSN: 0007-1188
- DT Article
- LA English
- L6 ANSWER 29 OF 45 BIOSIS COPYRIGHT 2000 BIOSIS
- AN 1993:413296 BIOSIS
- DN PREV199396079021
- TI Nafamostat mesilate: A regional anticoagulant for hemodialysis in patients
  - at high risk for bleeding.
- AU Akizawa, Tadao (1); Koshikawa, Shozo; Ota, Kazuo; Kazama, Mutsuyoshi; Mimura, Nobuhide; Hirasawa, Yoshihei
- CS (1) Dep. Internal Med., Showa Univ., Fujigaoka Hosp., 1-30 Fujigaoka, Midori-ku, Yokohama 227 Japan
- SO Nephron, (1993) Vol. 64, No. 3, pp. 376-381. ISSN: 0028-2766.
- DT Article
- LA English
- L6 ANSWER 30 OF 45 BIOSIS COPYRIGHT 2000 BIOSIS
- AN 1993:220149 BIOSIS
- DN PREV199344104649
- TI Intravenous FUT-175 inhibits complement activation in the cerebrospinal fluid and vasospasm-related delayed ischemic neurological deficit following subarachnoid hemorrhage.
- AU Yanamoto, H. (1); Kikuchi, H. (1); Okamoto, S.; Ishikawa, J.; Matsumoto, M.; Shimizu, Y.; Sato, M.; Tokuriki, Y.; Matsumoto, K.; Nakamura, M.
- CS (1) Dep. Neurosurg., Kyoto Univ. Med. Sch., Kyoto Japan
- SO Canadian Journal of Neurological Sciences, (1993) Vol. 20, No. SUPPL. 1, pp. S30.

  Meeting Info.: Vth International Symposium on Cerebral Vasospasm Edmonton and Jasper, Alberta, Canada May 18-21, 1993
  ISSN: 0317-1671.
- DT Conference
- LA. English
- L6 ANSWER 31 OF 45 CAPLUS COPYRIGHT 2000 ACS
- AN 1993:404342 CAPLUS
- DN 119:4342
- TI Efficacy of futhan rinse solution following rat heart preservation
- AU Urushihara, Takashi; Sumimoto, Kazuo; Sumimoto, Ryo; Ikeda, Masanobu; Fukuda, Yasuhiko; Dohi, Kiyohiko
- CS Sch. Med., Hiroshima Univ., Hiroshima, Japan
- SO Nippon Geka Gakkai Zasshi (1992), 93(12), 1514 CODEN: NGGZAK; ISSN: 0301-4894
- DT Journal
- LA Japanese
- L6 ANSWER 32 OF 45 BIOSIS COPYRIGHT 2000 BIOSIS
- AN 1993:288293 BIOSIS
- DN PREV199345006418
- TI Nafamostat mesilate saves the blood loss during open heart surgery.
- AU Murase, Mitsuya; Usui, Akihiko; Maeda, Masanobu; Tomita, Yasuhiro; Murakami, Fumihiko; Teranishi, Katuhito; Koyama, Tomio; Abe, Toshio
- CS Nagoya Univ., Nagoya, Aichi Japan
- SO Circulation, (1992) Vol. 86, No. 4 SUPPL. 1, pp. I567.
  Meeting Info.: 65th Scientific Sessions of the American Heart Association
  New Orleans, Louisiana, USA November 16-19, 1992
  ISSN: 0009-7322.
- DT Conference
- LA English
- L6 ANSWER 33 OF 45 BIOSIS COPYRIGHT 2000 BIOSIS
- AN 1992:263628 BIOSIS

- DN BA93:139953 EXPERIMENTAL STU IN A RABBIT MODEL OF ISCHEMIA-PERFUSION LUNG INJURY ΤI DURING CARDIOPULMONARY BYPASS. ΑU KURATANI T; MATSUDA H; SAWA Y; KANEKO M; NAKANO S; KAWASHIMA Y CS FIRST DEP. SURG., OSAKA UNIV. MED. SCH., 1-1-50 FUKUSHIMA, FUKUSHIMA-KU, OSAKA 553, JAPAN. SO J THORAC CARDIOVASC SURG, (1992) 103 (3), 564-568. CODEN: JTCSAQ. ISSN: 0022-5223. FS BA; OLD LAEnglish ANSWER 34 OF 45 CAPLUS COPYRIGHT 2000 ACS L6 DUPLICATE 2 1992:248072 CAPLUS AN 116:248072 DN TIEffect of anticomplement reagents, K-76 COOH and FUT175, on discordant xenograft survival Miyagawa, S.; Shirakura, R.; Matsumiya, G.; Kitagawa, S.; Fukushima, N.; ΑU Nakata, S.; Nakano, S.; Kitamura, H.; Matsumoto, M.; et al. Med. Sch., Osaka Univ., Osaka, Japan CS Transplant. Proc. (1992), 24(2), 483-4 SO CODEN: TRPPA8; ISSN: 0041-1345 DTJournal English LAL6ANSWER 35 OF 45 BIOSIS COPYRIGHT 2000 BIOSIS 1992:215286 BIOSIS ΑN DN BA93:115511 THERAPEUTIC TRIAL OF CEREBRAL VASOSPASM WITH THE SERINE PROTEASE TIINHIBITOR FUT-175 ADMINISTERED IN THE ACUTE STAGE AFTER SUBARACHNOID HEMORRHAGE. YANAMOTO H; KIKUCHI H; SATO M; SHIMIZU Y; YONEDA S; OKAMOTO S ΑU CS DEP. NEUROSURGERY, KYOTO UNIV. MED. SCH., KAWAHARA-CHO 54, SYOGOIN, SAKYO-KU, KYOTO, JPN. SO NEUROSURGERY (BALTIMORE), (1992) 30 (3), 358-363. CODEN: NRSRDY. FS BA; OLD English LAANSWER 36 OF 45 CAPLUS COPYRIGHT 2000 ACS L6 DUPLICATE 3 1991:526765 CAPLUS ΑN 115:126765 DN Beneficial effect of therapeutic infusion of nafamostat mesilate ΤI
- TI Beneficial effect of therapeutic infusion of nafamostat mesilate (FUT-175)

on hemodynamics in experimental acute pancreatitis

- AU Dobosz, M.; Sledzinski, Z.; Juszkiewicz, P.; Babicki, A.; Stanek, A.; Wajda, Z.; Basinski, A.
- CS 2nd Dep. Gen. Surg., Med. Acad., Gdansk, Pol.
- SO Hepato-Gastroenterology (1991), 38(2), 139-42 CODEN: HEGAD4; ISSN: 0172-6390
- DT Journal
- LA English
- L6 ANSWER 37 OF 45 CAPLUS COPYRIGHT 2000 ACS
- AN 1992:604810 CAPLUS
- DN 117:204810
- TI Combined administration of protease inhibitor and thromboxane A2 synthetase inhibitor for anticoagulation of a left ventricular assist device
- AU Takahama, Tatsuhiko; Kanai, Fukuei; Hiraishi, Mamoru; Onishi, Kiyoshi; Yamazaki, Zenya; Naruse, Yoshihiro; Furuse, Akira; Yoshitake, Tsuyoshi
- CS Saitama Med. Cent., Saitama Med. Coll., Kawagoe, 350, Japan
- SO ASAIO Trans. (1990), 36(3), M141-M144 CODEN: ASATEJ; ISSN: 0889-7190
- DT Journal
- LA English

- ANSWER 38 OF 45 L6 OSIS COPYRIGHT 2000 BIOSIS ΑN 1990:418614 BIO . DN BA90:79415 ΤI EFFECT OF NAFAMOSTAT MESILATE ON SERUM ACTIVITIES OF PANCREATIC ENZYMES AND PLASMA HORMONE LEVELS. ΑU WAKAYAMA S; SUZUKI T; SAKAI T; MATSUKI A CS DEP. OF ANESTHESIA, AOMORI ROSAI HOSP., HACHINOHE 031, JPN. SO JPN J ANESTHESIOL, (1990) 39 (6), 734-740. CODEN: MASUAC. ISSN: 0021-4892. FS BA: OLD LA Japanese L6 ANSWER 39 OF 45 BIOSIS COPYRIGHT 2000 BIOSIS 1991:20572 BIOSIS ΑN DN BR40:8902 COMPLEMENT ACTIVATION IN THE ISOLATED HEART PROTECTION BY ΤI FUT-175 NAFAMOSTAT. HOMEISTER J W; SATOH P S; LUCCHESI B R ΑU UNIV: MICH. MED. SCH., ANN ARBOR, MICH. CS SO 63RD SCIENTIFIC SESSIONS OF THE AMERICAN HEART ASSOCIATION, DALLAS, TEXAS, USA, NOVEMBER 12-15, 1990. CIRCULATION. (1990) 82 (4 SUPPL 3), III148. CODEN: CIRCAZ. ISSN: 0009-7322. DT Conference FS BR; OLD LA English L6 ANSWER 40 OF 45 BIOSIS COPYRIGHT 2000 BIOSIS 1989:350455 BIOSIS ΑN BR37:41552 DN PREVENTION OF PULMONARY EDEMA IN AUTOPERFUSING HEART-LUNG ΤI PREPARATION BY FUT-175 AND LEUKOCYTE DEPLETION. NAKA Y; HIROSE H; MATSUDA H; NAKANO S; SHIRAKURA R; KAWAGUCHI A; MIYAMOTO ΑU Y; MIYAGAWA S; FUKUSHIMA N; KAWASHIMA Y OSAKA UNIV. MED. SCH., FIRST DEP. SURGERY, 1-1-50, FUKUSHIMA, CS FUKUSHIMA-KU, OSAKA 553, JPN. TWELFTH INTERNATIONAL CONGRESS OF THE TRANSPLANTATION SOCIETY, SYDNEY, SO NEW SOUTH WALES, AUSTRALIA, AUGUST 14-19, 1988. TRANSPLANT PROC. (1989) 21 (1 PART 2), 1353-1356. CODEN: TRPPA8. ISSN: 0041-1345. BR; OLD FS English LA ANSWER 41 OF 45 CAPLUS COPYRIGHT 2000 ACS L6 1989:546489 CAPLUS ΑN 111:146489 DN Experimental study on the usefulness of the protease inhibitor, ΤI nafamostat mesilate (FUT), as an anticoagulant in left heart bypass ΑU Saito, A.; Moro, H.; Eguchi, S.; Yokosawa, T. Sch. Med., Niigata Univ., Niigata, Japan CS Jinko Zoki (1989), 18(2), 453-6 SO CODEN: JNZKA7; ISSN: 0300-0818 DTJournal Japanese LA ANSWER 42 OF 45 CAPLUS COPYRIGHT 2000 ACS L6 1988:31609 CAPLUS ΑN 108:31609 DN TΙ Comparative study of anticoagulation therapy with an LVAD system AU
- AU Takahama, Tatsuhiko; Kanai, Fukuei; Hiraishi, Mamoru; Onishi, Kiyoshi; Yamazaki, Zenya; Furuse, Akira; Asano, Kenichi; Yoshitake, Tsuyoshi
  CS Saitama Med. Cent., Saitama Med. Coll., Kawagoe, 350, Japan
  SO ASAIO Trans. (1987), 33(3), 227-34
- SO ASAIO Trans. (1987), 33(3), 227-34 CODEN: ASATEJ; ISSN: 0889-7190

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DT
     Journal
     English
LA
     ANSWER 43 OF 45 BIOSIS COPYRIGHT 2000 BIOSIS
     1986:16656 BIOSIS
AN
     BR30:16656
DN
     COMPLEMENT ACTIVATION DURING EXPERIMENTAL CARDIOPULMONARY BYPASS
TI
     AND INHIBITORY EFFECT OF FUT-175.
ΑU
     MIYAMOTO Y; HIROSE H; MATSUDA H; NAKANO S; SASAKO Y; NISHIGAKI K; TAKAMI
     H; KAWASHIMA Y; KITAMURA H; NAGAKI K
     22ND ANNUAL MEETING OF THE JAPANESE SOCIETY FOR ARTÍFICIAL ORGANS AND
SO
     TISSUES, OSAKA, JAPAN, NOV. 9-10, 1984. ARTIF QRGANS. (1985) 9 (3), 310.
     CODEN: ARORD7. ISSN: 0160 564X.
DT
     Conference
FS
     BR; OLD
     English
LA
                              COPYRIGHT 2000 BIOSIS
L6
     ANSWER 44 OF 45 BIOSIS
     1985:235770 BIOSIS
ΑN
     BA79:15766
DN
     PHARMACOLOGICAL STUDIES OF FUT-17 NAFAMSTAT MESILATE 1. INHIBITION OF
ΤI
     PROTEASE ACTIVITY IN-VITRO AND IN-VIVO EXPERIMENTS.
     AOYAMA T; INO Y; OZEKI M; ODA M; SATO T; KOSHIYAMA Y; SUZUKI S; FUJITA M
ΑU
     RES. LAB., TORII CO., LTD., 3-14-3 MINAMIYAWATA, ICHIKAWA, CHIBA 272,
CS
     JAPAN.
     JPN J PHARMACOL, (1984) 35 (3), 203-228. CODEN: JJPAAZ. ISSN: 0021-5198.
SO
FS
     BA; OLD
     English
LA
     ANSWER 45 OF 45 BIOSIS COPYRIGHT 2000 BIOSIS
L6
     1982:267691 BIOSIS
AN
DN
     BA74:40171
     EFFECTS OF FUT-175 IN ENDO TOXIN SHOCK.
ΤI
     EBATA T; KOBAYASHI K; DENNO R; GOTOH Y; AZUMA K; TOTSUKA M; HAYASAKA H
ΑU
     FIRST DEP. SURGERY, SAPPORO MED. COLLEGE, SAPPORO, 060.
CS
     JPN J ANESTHESIOL, (1982) 31 (1), 56-61.
SO
     CODEN: MASUAC. ISSN: 0021-4892.
FS
     BA; OLD
LA
     Japanese
=> d 6, 8, 9, 13, 16 ab, bib
     ANSWER 6 OF 45 CAPLUS COPYRIGHT 2000 ACS
L6
     It is well known that activation of proteases in the lysosomes and
AΒ
cytosol
     is one of the mechanisms of ischemic injury. It might thus be beneficial
     to det. whether the addn. of several clin. available protease inhibitors
     to a cardioplegic soln. can improve its protective ability.
     Using an isolated working rat heart prepn., the effects of
     several protease inhibitors (serine protease inhibitors; nafamostat
     mesylate and gabexate mesylate, a thiol-protease inhibitor; NCO-700; and
а
     urinary trypsin inhibitor, urinastatin) on the postischemic recovery of
     function and enzyme leakage were investigated in this study. These
     protease inhibitors were added to either the cardioplegic soln.
     or reperfusion soln. The addn. of each of the protease inhibitors,
except
     urinastatin, to the cardioplegic soln. improved the postischemic
     recovery of function and reduced enzyme leakage. The dose-response
     characteristics of these three protease inhibitors were bell shaped, and
     the optimal concns. of nafamostat mesylate, gabexate mesylate, and
```

NCO-700 were 5 .mu.M, 100 .mu.M, and 20 .mu.M, resp. In contrast to the results

of the preisches treatment study, the addn. of y of the protease inhibitors to the perfusion medium during Langence of reperfusion failed to improve the postischemic recovery of function and to reduce enzyme leakage. Surprisingly, the addn. of NCO-700 to the reperfusion soln. at

а

concn. of 5 .mu.M or higher had rather harmful effects on both functional recovery and enzyme leakage. These findings suggest that serine and

proteases may play an important role in myocardial injury during ischemia,

but not necessarily during reperfusion.

ΑN 1998:183 CAPLUS

DN 128:110669

Effects of protease inhibitors on postischemic recovery of the TΙ

Shibata, Toshihiko; Yamamoto, ΑU Fumio; Suehiro, Shigefumi; Kinoshita,

Second Dep. of Surgery, Osaka Lity University Medical School, Osaka, 545, CS Japan

Cardiovasc. Drugs Ther. (1997) / 11(4), 547-556 SO CODEN: CDTHET; ISSN: 0920-3206,

Kluwer Academic Publishers PB

DT Journal

English LA

ANSWER 8 OF 45 BIOSIS COPYRIGHT 2000 BIOSIS L6

New therapies of cerebral vasospasm aim to prevent the effects of AΒ subarachnoid haemorrhage. These effects result in red blood cell haemolysis and release of oxyhaemoglobin, free radicals formation and lipid peroxidations, imbalance in endothelial modulation of vasomotor

tone

and activation of the complement system. Low doses of fibrinolytic agents administered intrathecally accelerate the fibrinolysis of the clot and reduce the oxyhaemoglobin release. The tissue-type plasminogen activator has proven to be effective in preventing vasospasm, but the modalities of this therapy remain to be defined. Free radical reactions may be inhibited

by free radical scavengers and inhibitors of lipid peroxidabons. Tirilazad

is a potent inhibitor of lipid peroxidatons, which improves the patients' outcome and has gone to Phase III human trials. Superoxide dismutase and tropolone derivatives are currently evaluated in animal models. Vasomotor tone can be modified in experimental models either by blocking endothelin receptors (BQ-123), or by facilitating the release and enhancing the effect of nitric oxide using protein kinase C inhibitors, drugs that increase intracellular calcium (cyclopiazonic acid, LP-805) and free radicals scavengers (superoxide dismutase). These possibilities are being investigated. Finally preliminary studies have demonstrated the efficacy of FUT-175, an inhibitor of the complement system, in the prevention of vasospasm. In the next years, these new therapies have to be validated by prospective and randomized clinical trials to propose guidelines for the management of patients at risk of cerebral vasospasm after aneurysmal rupture.

1996:321861 BIOSIS AN

DN PREV199699044217

Pharmacological therapeutic prospects of cerebral vasospasm. TΙ

ΑU Hans, P.

Serv. Univ. d'Anesthesie Reanim., CHR de la Citadelle, 4000 Liege Belgium CS

Annales Francaises d'Anesthesie et de Reanimation, (1996) Vol. 15, No. 3, SO pp. 374-381. ISSN: 0750-7658.

 $\mathsf{DT}$ General Review

LA French

SL French; English

ANSWER 9 OF 45 CAPLUS COPYRIGHT 2000 ACS L6

The effects of amostat mesilate (NM) on myoca al, biochem., and functional changes in canine hearts were examd. In isolated heart was preserved for 6 h at 5.degree. and then reperfused for 2 h at 37.degree. NM was added to the cardioplegic soln. At both 10-7M and 10-6M, NM was able to maintain myocardial cAMP at a normal level and to reduce cGMP concres. at the end of both preservation and reperfusion. The serum N-acetyl-.beta.-D-glucosaminidase concn. during reperfusion was lower in hearts treated with NM 10-6 or 10-7M NM than in those without NM. Although NM failed to preserve myocardial concns. of adenine nucleotide compds., NM at 10-7M maintained the .+-. dp/dt of the left ventricle after reperfusion at the same level as in the nonischemic control group and better than NM at 10-6M or no NM. Myocardial uptake of 10-5M NM was 55% during the 6-h preservation and 29% during the 2-h reperfusion. It is concluded that addn. of 10-7M NM to a nondepolarizing soln. does not preserve myocardial adenine nucleotide concns. but does facilitate the recovery of left ventricular function.

NM

at 10-5M seems to have a high affinity for the myocardium and may depress the recovery of left ventricular function.

AN 1997:31579 CAPLUS

DN 126:84283

TI The effect and pharmacokinetics of nafamostat mesilate adjunct to cold nondepolarizing cardioplegia in a canine model of cardiac preservation

AU Sunamori, Makoto; Yoshida, Tetsuya; Miyamoto, Hisashi; Wang, Yigang; Suzuki, Akio

CS School Medicine, Tokyo Medical and Dental University, Tokyo, 113, Japan

SO Transplant Int. (1996), 9 (4), 364-369 CODEN: TRINE5; ISSN: 0934-0874

PB Springer

DT Journal

LA English

L6 ANSWER 13 OF 45 BIOSIS COPYRIGHT 2000 BIOSIS

 ${\tt AB}$  In cardiac operations endopeptidase (protease) inhibitor may be beneficial

in reducing myocardial injury when administered in the <code>cardiopulmonary</code> bypass prime. Nafamostat mesilate was evaluated in 20 patients who underwent coronary artery bypass grafting. The patients were divided into a control group (n=10) and a nafamostat group (n=10). Nafamostat (2 mg/kg per hour) was continuously given during <code>cardiopulmonary</code> bypass in the nafamostat group. The age, number of grafts, <code>cardiopulmonary</code> bypass time, and aortic crossclamp time were similar between groups. In the control group, neither tumor necrosis factor-alpha nor interleukin-1 levels showed any significant change

cardiopulmonary bypass, whereas interleukin-6 and interleukin-8
levels, percent expression of adhesion molecule (CD18) on neutrophils,
and

CH-50 assay results increased significantly during cardiopulmonary bypass. As compared with the control group, the nafamostat group showed significantly lower levels of interleukin-6 (123 +- 57 versus 40 +- 22 pg/ml, respectively,) and interleukin-8 (96 +- 13 versus 66 +- 14 pg/ml, respectively). The nafamostat group showed a significantly lower difference of CH-50 assay results and malondialdehyde levels between coronary sinus blood and arterial blood and peak values of creatine kinase

MB (43 +- 12 IU/L versus 19 +- 6 IU/L) during the postoperative course compared with findings in the control group. These results demonstrated that inflammatory reactions induced by  ${\bf cardiopulmonary}$  bypass had adverse effects on myocardial recovery after aortic crossclamping and that nafamostat mesilate given during  ${\bf cardiopulmonary}$  bypass appeared to reduce myocardial reperfusion injury by attenuating such inflammatory reactions. Attenuation of inflammatory reactions of  ${\bf cardiopulmonary}$  bypass should be considered in the strategy of myocardial protection.

- AN1996:112706 BIG PREV19969868484 DN
- TΙ Attenuation of cardiopulmonary bypass-derived inflammatory
- reactions reduces myocardial reperfusion injury in cardiac operations. ΑU Sawa, Yoshiki; Shimazaki, Yasuhisa; Kadoba, Keishi; Masai, Takashi; Fukuda, Hirotsugu; Ohata, Toshihiro; Taniguchi, Kazuhiro; Matsuda, Hikaru
- CS (1) First Dep. Surgery, Osaka Univ. Med. Sch., 2-2 Yamada-oka, Suita, Osaka 565 Japan
- Journal of Thoracic and Cardiovascular Surgery, (1996) Vol. 111, No. 1, SO pp. 29-35. ISSN: 0022-5223.
- DT Article
- English LA
- ANSWER 16 OF 45 BIOSIS COPYRIGHT 2000 BIOSIS L6
- 1995:243942 BIOSIS AN
- PREV199598258242 DN
- TΙ The anti-complement effects of FUT-175 on myocardial ischemia/reperfusion injury in the blood-perfused isolated rabbit hearts.
- Yokota, Syunji (1); Kan-No, Satoshi; Saitoh, Yoshiaki; Kasama, Kikuko; Ohara, Naoki; Ono, Hiroshi
- CS (1) Lab. Applied Pharmacology, Hatano Res. Inst., Food Drug Safety Center,

Kanagawa 257 Japan

SO Japanese Journal of Pharmacology, (1995) Vol. 67, No. SUPPL. 1, pp. 280P.

Meeting Info.: 68th Annual Meeting of the Japanese Pharmacological Society

Nagoya, Japan March 25-28, 1995

ISSN: 0021-5198.

DT Conference

LA English

- => s myocardial infarction? or stroke? or hemorrhagic shock? or diabectic retinopathy? or venous insufficiency?
- L7399340 MYOCARDIAL INFARCTION? OR STROKE? OR HEMORRHAGIC SHOCK? OR DIABE

CTIC RETINOPATHY? OR VENOUS INSUFFICIENCY?

- => s diabetes?
  - 3 FILES SEARCHED...
- $\Gamma8$ 320264 DIABETES?
- => s 17 or 18
- 707481 L7 OR L8
- => s 19 and 11
- 6 L9 AND L1 L10
- => d 1-6 ab,bib
- ANSWER 1 OF 6 CAPLUS COPYRIGHT 2000 ACS L10
- Diagnostic methods that rely on the use of one or more assays that assess AΒ cellular attivation are provided. The assays are performed on whole
  - or leuko tes (neutrophils), and indicate individually or in combination the level of cardiovascular cell activation, which is pivotal in many chronic and acute disease states. These results of the assays are used within a clin. framework to support therapeutic decisions such as:

further

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testing for inf jous agents, anti-oxidant or a adhesion therapy, postponement and ptimal re-scheduling of high-risk surgeries,
classifying
     susceptibility to and progression rates of chronic disease such as
     diabetes, organ rejection, atherogenesis, and venous
     insufficiency; extreme interventions in trauma cases of
     particularly high risk and activation-lowering therapies. Also provided
     is compn. derived from a pancreatic homogenate that contains circulating
     cell activating factors, which can serve as targets for drug screening to
     identify drug candidates for use in activation lowering therapies.
     Methods for lowering cell activation by administering protease
inhibitors,
     particularly serine protease inhibitors, are also provided. Kits for
     performing the methods are also provided.
     1999:595348 CAPLUS
     131:225828
     Methods of diagnosis and triage using cell activation measures
     Stoughton, Roland B.; Schmid-Schonbein, Geert W.; Hugli, Tony E.;
IN
Kistler,
     Erik
     Cell Activation, Inc., USA; The Regents of the University of California;
     The Scripps Research Institute
     PCT Int. Appl., 184 pp.
SO
     CODEN: PIXXD2
DT
     Patent
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     English
FAN.CNT 1
     PATENT NO.
                     KIND DATE
                                             APPLICATION NO. DATE
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     WO 9946367
                       A2
                             19990916
                                             WO 1999-US5247 19990311
PΙ
                      A3 19991209
     WO 9946367
         W: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ,
             DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS,
             JP, KE, KG, KP, KR, KZ, LC, LR, LS, LT, LU, LV, MD, MG, MK, MN,
             MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZW, AM, AZ, BY, KG, KZ, MD, RU,
             TJ, TM
         RW: GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW, AT, BE, CH, CY, DE, DK,
             ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
     AU 9931829
                       A1 19990927
                                            AU 1999-31829
                                                               19990311
PRAI US 1998-38894
                       19980311
     WO 1999-U$5247
                       19990311
L10
     ANSWER 2 OF 6 CAPLUS COPYRIGHT 2000 ACS
AΒ
     Hypercoagulability is known to occur in the early phase of
     hemorrhagic shock. The prolongation of excessive clot
     formation after recovery from a shock state leads to the formation of
     microthrombi or disseminated intravascular coagulation which disturbs
     microcirculation, damaging organ function. The aim of the present study
     is to investigate the beneficial effect of a synthetic protease
inhibitor,
     6-amidino-2-naphthyl p-guanidinobenzoate dimethanesulfonate (nafamostat
     mesilate), in the attenuation of hypercoagulability in hemorrhagic
     shock. A model of hemorrhagic shock that
     simulates the clin. course of injured patients was created in
anesthetized
     dogs. The animals were divided into 2 groups: a control group (group-C)
     and an exptl. group (group-E). Animals received saline or 0.2 mg/kg of
     nafamostat mesilate resp., when their mean arterial pressure declined to
     50 mmHg. The serum concn. of hydroxytryptamine (5-HT), prothrombin time
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(PT), and activated partial thromboplastin time (APTT) were detd. as indicators of platelet activity and blood coagulation. In group-C, serum 5-HT was elevated significantly at 60 min after hemorrhagic shock but not so in group-E. The APTT at 30 and 60 min was shorter in group-C than in group-E. The PT at 30 min was also shorter in

group-C. Plasm birin degrdn. products (FDP) is eased at 60 min after the induction of nock in group-C. The results indicate that inadequate tissue perfusion in shock stimulates blood coagulation and that mesilate might be beneficial in decreasing excessive blood coagulation. ΑN 1997:186530 CAPLUS DN TINafamostat mesilate, a synthetic protease inhibitor, attenuated hypercoagulability in a canine model of hemorrhagic Koido, Yuichi; Kato, Kazuyoshi; Shimizu-Suganuma, Masumi; Shichinohe, ΑU Kazuhiro . Dep. Emerg. Crit. Care Med., Nippon Med. Sch., Tokyo, 113, Japan Nippon Ika Daigaku Zasshi (1997), 64(1), 9-15 CODEN: NIDZAJ; ISSN: 0048-0444 PΒ Nippon Ika Daigaku Igakkai DTJournal LA English L10 ANSWER 3 OF 6 CAPLUS COPYRIGHT 2000 ACS Acute pancreatitis was induced in 13 anesthetized dogs by retrograde. injection of bile mixed with trypsin into the pancreatic duct. Six animals were treated with i.v. infusion of new synthetic antiprotease, Nafamostat Mesilate, at a dose of 1 mg/kg/h. Four out of 7 untreated animals died during the expt. All the treated dogs survived. Hemodynamic data were regularly monitored during a 10-h observation period. Cardiac output, mean arterial pressure and left ventricular stroke vol. decreased rapidly in the untreated animals. An increase in systemic vascular resistance and pulmonary vascular resistance was obsd. in dogs without treatment. Nafamostat Mesilate given as therapy significantly improved the hemodynamic parameters, and prevented the animals from developing shock. The study demonstrates an advantageous influence of synthetic antiprotease Nafamostat Mesilate on the course of acute exptl. pancreatitis. AN1991:526765 CAPLUS 115:126765 DN ΤI Beneficial effect of therapeutic infusion of nafamostat mesilate (FUT-175) on hemodynamics in experimental acute pancreatitis Dobosz, M.; Sledzinski, Z.; Juszkiewicz, P.; Babicki, A.; Stanek, A.; ΑU Wajda, Z.; Basinski, A. 2nd Dep. Gen. Surg., Med. Acad., Gdansk, Pol. Hepato-Gastroenterology (1991), 38(2), 139-42 CODEN: HEGAD4; ISSN: 0172-6390 CS SO DT Journal English LAANSWER 4 OF 6 BIOSIS COPYRIGHT 2000 BIOSIS Background: Reperfusion injury in the myocardium has recently been considered to be a type of inflammation, and close attention has been to the possible involvement of neutroph/ls, complement, and cytokines in the onset of this injury. Recently, it has been reported that serum levels of interleukin-6 are elevated significantly after myocardial infarction. The major site of interleukin-6 production and its exact roles are still unknown. In this study, we hypothesized that myocytes may produce interleukin-6-during hypoxia and this may play a in neutrophil-mediated reperfusion injury. Methods and results: In the

clinical study, 20 patients who underwent coronary artery bypass grafting were divided into 2 groups: group F, in which patients were treated with

serine protease inhibitor (FUT-175, 2 mg/kg per hour) during cardiopulmonary bypass, and group C (untreated patients). In group C,

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myocardial interpukin-6 production, as determined by the difference between the interpulsion of level in the cardiopuls. By by ass circuit and its level in coronary venous blood, increased significantly after reperfusion (12 +- 4 pg/mL) as compared with that before aortic crossclamping (2 +- 2 pg/mL). In group F, the increase in the interleukin-6 level was suppressed significantly (before aortic crossclamping, 3 +- 2 pg/mL; after reperfusion, 4 +- 3 pg/mL). The interleukin-6 production differed significantly between group C and group F. In the in vitro experimental study, the supernatant from myocytes exposed to 2 hours of hypoxia (group 2H) showed significantly higher levels of interleukin-6 (455 +- 260 pg/mL) than that from normoxic myocytes (group N) (47 +- 15 pg/mL). This interleukin-6 production was suppressed by the addition of FUT-175 (123 +- 24 pg/mL). The interleukin-6

 $\hbox{production by endothelial cells of coronary vessels did not differ}\\$ 

group 2H (283 +- 151 pg/mL) and group N (151 +- 86 pg/mL). In a coincubation system with a monolayer of endothelial cells on collagen membrane and myocytes under collagen membrane in a modified Boyden chamber, 2 hours of coincubation showed a significantly higher percent of neutrophil transendothelial migration (group 2H vs N, 78% +- 13% vs 26%

11%), value of chemiluminescence (22 +- 8 vs 5 +- 2 X 103 counts/3 minutes), and percent of irreversibly damaged myocytes (48% +- 17% vs 12% +- 8%) than normoxic coincubation. In contrast, anti-interleukin-6 monoclonal antibody significantly attenuated neutrophil transendothelial migration (42% +- 19%) and irreversible damage of myocytes (26% +- 15%)

2 hours of coincubation. Conclusions: Interleukin-6 is produced from myocardium during ischemia and reperfusion in patients undergoing coronary

bypass grafting. This interleukin-6 may be derived from hypoxic myocytes and play a role in neutrophil-mediated reperfusion injury in myocardium.

AN 1998:475040 BIOSIS

DN PREV199800475040

TI Interleukin-6 derived from hypoxic myocytes promotes neutrophil-mediated reperfusion injury in myocardium.

AU Sawa, Yoshiki (1); Ichikawa, Hajime; Kagisaki, Koji; Ohata, Toshihiro; Matsuda, Hikaru

CS (1) First Dep. Surg., Osaka Univ. Med. Sch., 2-2 Yamada oka, Suita, Osaka 565 Japan

SO Journal of Thoracic and Cardiovascular Surgery, (Sept., 1998) Vol. 116, No. 3, pp. 511-517. ISSN: 0022-5223.

DT Article LA English

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L10 ANSWER 5 OF 6 BIOSIS COPYRIGHT 2000 BIOSIS

AB A 41-year old male with insulin-dependent diabetes mellitus previously unsuccessfully treated with a controlled diet and glibenclamide, and subsequently with increasing insulin doses (5 and 20 IU/day) experienced polyuria, glycosuria and loss of weight. On admittance

to hospital serum C3 concentrations were found to be depressed. The insulin dose was further increased to 30 IU/day and the patient was also treated with 20 mg nafamostat mesylate given intravenously twice daily

 $6~{\rm days}.$  On completion of nafamostat mesylate treatment serum C3 concentrations were increased but after 17 days they started to decrease again.

AN 1991:453761 BIOSIS

DN BA92:98541

TI COMPLEMENT ACTIVATION VIA THE ALTERNATIVE PATHWAY IN A PATIENT WITH INSULIN-DEPENDENT DIABETES MELLITUS.

AU OKADA S; ICHIKI K; TANOKUCHI S; ISHII K; HAMADA H; YAMAMOTO H; OTA Z CS THIRD DEP. MED., OKAYAMA UNIV. MED. SCH., 2-5-1 SHIKATA-CHO, OKAYAMA 700,

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JPN.

SO J INT MED RES, 91) 19 (4), 348-350. CODEN: JIMRBV. ISSN: 0300-0605.

FS BA; OLD

LA English

L10 ANSWER 6 OF 6 BIOSIS COPYRIGHT 2000 BIOSIS

AB Sera containing islet cell surface antibody were obtained from seven children with insulin-dependent diabetes mellitus soon after the onset of disease. After incubation of 51Cr-labelled rat islet cells with islet cell surface antibody, human AB-type serum with or without nafamostat mesylate was added before further incubation. Radioactivity in the supernatant was measured to determine complement-dependent antibody-mediated cytotoxicity. Cytotoxicity in untreated sera [mean]

SD) 19.4 .+-. 4.0%] was significantly (P < 0.001) inhibited by ethyleneglycoltetraacetic acid (EGTA) (7.1 .+-. 4.9%), ethylene diaminetetraacetic acid (EDTA) (2.5 .+-. 0.9%) and nafamostat mesylate (2.8 .+-. 1.8%). Cytotoxicity of nafamostat mesylate-treated serum was significantly (P < 0.05) lower than that of EGTA-treated serum but not significantly different from that of EDTA-treated serum. There was no difference in cytotoxicity between nafamostat mesylate-treated and untreated, inactivated human serum. The results indicate that the otease

inhibitor nafamostat mesylate completely inhibited the complement activation of the immune complex associated with islet cell surface antibody by the classical and alternative pathways.

AN 1991:365062 BIOSIS

DN BA92:53287

TI DOES PROTEASE INHIBITOR INHIBIT COMPLEMENT ACTIVATION CAUSED BY THE IMMUNE

COMPLEX ASSOCIATED WITH ISLET CELL SURFACE ANTIBODY.

AU OKADA S; SATO K; ICHIKI K; TANOKUCHI S; ISHII K; OTA Z; TAKEDA A

CS THIRD DEP. MED. OKAYAMA UNIV. MED. SCH., OKAYAMA 700, JAPAN.

SO J INT MED RES, (1991) 19 (3), 234-236. CODEN: JIMRBV. ISSN: 0300-0605.

FS BA; OLD

LA English

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